

In the Specification:

Please replace Paragraph 0069 on pages 15-16 as follows:

[0067] In order to clone both complete toxin genes, a *Sau3A* partial library was constructed. PS81I total cellular DNA partially digested with *Sau3A* and size fractionated by electrophoresis into a mixture of 9-23 Kb fragments on a 0.6% agarose-TAE gel, and purified as described previously, was ligated into LambdaGEM-11TM (PROMEGA). The packaged phage were plated on P2392 *E. coli* cells (Stratagene) at a high titer and screened using the radiolabeled synthetic oligonucleotides (aforementioned) as nucleic acid hybridization probes. Hybridizing plaques, using each probe, were rescreened at a lower plaque density. Purified plaques that hybridized with either probe were used to infect P2392 *E. coli* cells in liquid culture for preparation of phage for DNA isolation. DNA was isolated by standard procedures. Preparative amounts of DNA were digested with *SalI* (to release the inserted DNA from lambda arms) and separated by electrophoresis on a 0.6% agarose-TAE gel. The large fragments, electroeluted and concentrated as described above, were ligated to *SalI*-digested and dephosphorylated pUC19 (NEB). The ligation mix was introduced by transformation into DH5(α) competent *E. coli* cells (BRL) and plated on LB agar containing ampicillin, isopropyl-(β)-D-thiogalactoside (IPTG), and 5-bromo-4-chloro-3-indolyl-(β)-D-galactoside (XGAL). White colonies, with prospective insertions in the (β)-galactosidase gene of pUC19, were subjected to standard rapid plasmid purification procedures to isolate the desired plasmids. Plasmid pM3,122-1 contains a 15 Kb *Sau3A* fragment isolated using the 81IA oligonucleotide probe. Plasmid pM4,59-1 contains an 18 Kb *Sau3A* fragment isolated using the 81IB oligonucleotide probe.